# COMPOSITION FOR COATING ORGANIC ELECTRODE AND METHOD OF MANUFACTURING ORGANIC ELECTRODE HAVING EXCELLENT TRANSPARENCY USING THE COMPOSITION

## 5 [Technical Field]

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The present invention relates to a composition for coating an organic electrode and method of manufacturing an organic electrode having an excellent transparency using the composition, and more particularly to a method of manufacturing an organic electrode having an excellent transparency comprising steps of mixing a aqueous solution of polyethylenedioxythiophene(PEDOT) conductive polymers having nano-sized particle,

polyhydric alcohol, polyol or a mixed solvent thereof, microphase-separating conductive polymers particle having a nano-sized from the aqueous solution of conductive polymers, wherein a visible ray transmittance of an organic conductive layer is more than 90% in case of coating and wherein a sheet resistance of layer is 300 to  $900\Omega/sq$ .

As computer, various household appliances and telecommunication device are digitalized and are rapidly highly effected, the embodiment of large layer and portable display have been necessarily needed. In order to embody a large portable flexible display, a display material which is possible to fold or roll such a newspaper has been needed.

To achieve the large portable flexible display, an electrode material using for display is transparent, and also shows a low resistance and high intensity so as to mechanically stabilized when being bent or folded. Further the electrode material should have a coefficient of thermal expansion similar to coefficient of thermal expansion of a plastic substrate so that a short circuit or large change of a sheet resistance should not occur in case of being overheated or in a high temperature.

It is possible to manufacture a display having any form by means of using a flexible display, therefore can be made the best use of clothes which can be changed a color or a pattern, trademark of clothes, billboard, price guidebook of commodity stand or a large area electrical light device as well as the portable display device.

30 [Background Art]

Patent Application

Presently, a study has been progressed about study about a chemical vapor deposition using a various metallic oxide of Indium, Tin, Zinc, Titanium and Cesium, magneton sputtering and a reactive evaporation as a method for preparing transparent electrode at home and abroad. But the study has a demerit resulting in high process cost in order to coat to a metal oxide on the substrate because coating metallic oxide to substrate needs to vacuum condition.

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As a method for preparing transparent electrode processed at lower cost, a method for using conductive polymers has been become influential. The electrode processed I by using conductive polymers is good for decreasing the process cost and working process because it is capable of using a various existing method for a coating polymer. In case of manufacturing a flexible display or a electrical light device, when the transparent electrode processed with the conductive polymers such a polyacethylene, polypyrrole, polyaniline and polythiophene is compared with the transparent electrode processed with a Indium Tin Oxide(ITO), it takes advantage of the process, and in case of need for a very flexible electrode, particularly manufacture of touch screen, the transparent electrode processed with the conductive polymers is good for increasing a life time. The transparent electrode have many merits, but generally conductive polymers increases a sheet resistance in case of slightly coating layer in order to increase the transmittance so that conductive polymers absorb a ray in the visible rays area and the conductive feature of organic electrode manufactured by the conductive polymers is increased in proportional to the thickness of the electrode. Therefore the conductive polymers is difficulty in applying to a applicable field of the transparent electrode such a touch panel, flexible display. Particularly in order to improve the process of the conductive polymers, when the transparent electrode is manufactured with the method, wherein the conductive polymers is pulverized to the nano-sized particle and uses commercially available dispersed polythiophene, it shows about  $1M\Omega/sq$  of sheet resistance. Therefore it is difficulty in using with the transparent electrode using for displaying in the condition of 85% of transmittance.

US 5,766,515, US6,083,635 and Korean Patent Publication No. 2000-1824 disclose a method for improving the conductivity of electrode manufactured with the aqueous solution of polyethylenedioxythiophene(PEDOT) conductive polymers having nano-sized particle by using a solvent or a additive. But US 5,766,515 or US6,083,635 has a problem as following, when a polyhydric alcohol, e.g sorbitol is added, the sheet resistance of coating layer having more than Patent Application 2 MJM Docket No. 4820-022

AJM Docket No. 4820-022 Client No. PPW05-076US 90% transmittance is difficult to decrease I less than  $\text{IM}\Omega/\text{sq}$ , and when a amide solvent is added, the sheet resistance of coating layer can be decrease less than  $\text{lk}\Omega/\text{sq}$ , but a hardness of layer is low and coating feature is diminished. On the other hand, according to Korean Patent Publication NO. 2000-1824, when a silicasol is added to aqueous solution of polythiopene treated with the amide solvent, the hardness of layer is improved, and the sheet resistance is increased more than  $\text{IM}\Omega/\text{sq}$  in case of improving the hardness of layer.

To consider above the feature, the coating layer shows more than 90% of the transmittance and less than several hundred  $\Omega$ /sq of the sheet resistance, also has the excellent transparency and hardness and the low resistance. Therefore a development of organic transparency electrode material which can apply to a electronic equipment has been needed continually.

[Disclosure]

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[Technical Problem]

Therefore, the present inventor has repeatedly studied a composition for coating an organic electrode for producing a organic electrode having a high transparency. Finally, the present inventors have found and completed that it makes to microphase-separate conductive polymers having nano-sized particle from a aqueous solution of conductive polymers when a aqueous solution of polyethylene dioxythiophene(PEDOT) conductive polymers having nano-sized particle, polyhydric alcohol and as a surfactant, a primary alcohol solvent and a amide solvent, a sulfoxide solvent or a mixed solvent thereof is mixed. Therefore in case of coating to the composition, the transmittance of conductive layer in the visible ray area shows more than 90% and the sheet resistance shows the range of 300 to 900Ω/sq.

An object of the present invention is to provide the composition for coating an organic electrode which can be microphage-separated the conductive polymers of nano-sized particle. Another object of the present invention is to provide a method for preparing high transparent organic electrode using the composition.

[Technical Solution]

To achieve the object of present invention, the feature of a composition for coating an organic electrode according to present invention is a composition for coating an organic electrode comprising 3% to 20% by weight of a polyhydric alcohol, a polyol or a mixture thereof; 5% to 10% by weight of a primary alcohol having C1 to C5; 5% to 25% by weight of a amide, sulfoxide or a mixed solvent thereof; 0. 01% to 0.1% by weight of a surfactant and an aqueous solution of polyethylene dioxythiophene (PEDOT) conductive polymers having nanosized particle in a remainder; and wherein a concentration of polyethylenedioxythiophene (PEDOT) and I polystyrenesulfonate (PSS) solid in the aqueous solution is 1.0% to 1.5% by weight of based on the total weight of solution, wherein a visible ray transmittance of organic conductive layer is more than 90% in case of coating, wherein a sheet resistance of layer is 300 to 900Ω/sq.

Also, the feature of the method for preparing high transparency organic electrode according to present invention is to comprise a method of preparing high transparent; organic electrode comprising steps of stirring the composition, spreading out the composition on a transparent substrate, drying up the substrate and coating 0.2 to 2.0µm by thickness of coating layer.

Also, the feature of a method for preparing high transparency organic electrode according to present invention is to comprise steps of stirring the composition, repeatedly dispersing the composition 2 to 10 times per 3 to 10minutes with a ultra I sonicator controlled by 20,000 to 40,000 Hz of frequency, 50 to 700W of power, spreading out the dispersed solution on the transparent substrate, drying up the substrate and coating 0.2 to 20μm by thickness of coating layer.

#### [Best Mode]

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Following, a composition for coating an organic electrode according to the present invention is detailedly described.

A composition for coating an organic electrode according to present invention is; comprising: as a essential constituent, an aqueous solution of polyethylenedioxythiophene(PEDOT) conductive polymers having nano-sized particle; a polyhydric alcohol, polyel or a mixture thereof; a primary alcohol having C 1 to C5; a amide, Patent Application 4 MJM Docket No. 4820-022

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sulfoxide or a mixed solvent thereof and a surfactant, and further comprising a dopant containing a crossing-linking agent or a sulfonic acid group(-SO<sub>3</sub>H).

The aqueous solution of polyethylenedioxythiophene(PEDOT) conductive polymers having nano-sized particle is dispersed some 5 repeating unit of a ethylene dioxythiophene oligomer into a polystyrenesulfonate(PSS) gel, wherein a concentration of the polyethylenedioxythiophene(PEDOT) and the polystyrene sulfonate(PSS) solid in aqueous solution is 1.0% to 1.5% by weight of based on the total weight of the aqueous solution, more preferably 0.4% to 0.7% by weight of the polyethylenedioxythiophene, 0.6% to 0.8% by weight of the polystyrene sulfonate(PSS). For example, as a aqueous solution of polyethylenedioxythiophene(PEDOT) conductive polymers having nano-sized particle, Baytron P(bayer Co.,Ltd) can be used in the present invention. On the other hand, the conductivity is not belong to the range of 300 to 900Ω/sq, when the aqueous solution of conductive polymers is less than 40% by weight of the composition for coating the; organic electrode.

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The transmittance of the visible ray areas is decreased less than 85% when the aqueous solution of conductive polymers is more than 70% by weight of a composition for coating organic electrode. Therefore, it is preferred that the conductive polymers do not belong to the range.

The polyhydric alcohol, polyol or the mixture thereof from among the constituent needs a affinity which is possible to mix with the nano particle of the conductive polymers in the metastable condition, simultaneously the function which is increased the conductivity among ethylenedioxythiophene by improving a cohesive force between the conductive nano particle by interaction with the polystyrene sulfonate(PSS) and the function which is improving the transmittance of the film by forming the empty space linked to each other conductive nano particle by the microphage-separation.

In order to improve a adhesive strength of each other particle, the polyhydric alcohol, polyol or the mixture thereof from among the constituent should contain more than two of hydroxy group(-OH). In order to simultaneously perform the adhesive strength between the conductive nano particle and the improvement of transmittance by microphage-separation, a molecule weight of polyhydric alcohol is preferable to less than 300. In case of more than 300m.w of polyhydric alcohol, a distance between the conductive nano particle become more Patent Application

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distance, therefore the conductivity may be; decreased. The example of usable alcohol is a ethyleneglycol, propylenegylcol, butanediol, neopentylgylcol, diethylenegylcol, triethylenegylcol, methylpentanediol, hexanediol, trimethylolpropane, glycerine, ethylhexanediol, hexanetriol, polyethyleneglycol, polypropyleneglycol, polyoxypropyleneglycol, polytetram ethyleneglycol, sorbitol and a derivative thereof, more preferably a ethyleneglycol, di ethylenegylcol or glycerine of less than 150m.w. A improvement of conductivity by the additive and the hardness of layer is not effective when the polyhyric alcohol or polyol is less than 3% by weight. The conductivity is lowered by relatively decreasing! the weight of nano particle of conductivity polymer when the polyhyric alcohol or the polyol is less than 20% by weight. Therefore, the polyhydric alcohol, polyol or the mixture thereof is preferable to use 3% to 20% by weight of based on the total weight of the composition for coating organic electrode.

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On the other hand, the amide solvent and sulfoxide solvent from among the constituent of the present invention easily make to swell a gel to be superior to the affinity with the polystyrenesulfonate(PSS) as a dopant forming nano particle gel of conductive polymers. The conductive nano particle is formed a bend and is easy to percolate between dispersed ethylenedioxythiophene oligomer by means of interactive diffusion of a chain of polymers between swelling gel, therefore the conductivity is improved. For example, as a amide solvent, a formamide, N-methylformamide, N,N-dimethylformamide, acetamide, N-methylacetamide, N,N-dimethylacetamide, N-methylpropionamide, 2- pyrrolidone, N-methylpyrrolidone, caprolactam and a 1,1,3,3- tetramethylurea can be used. Also as a sulfoxide solvent, a methylsulfoxide, dimethylsulfoxide, sulfolane and a diphenylsulfone can be used. When the amide, sulfoxide or the mixed solvent thereof is less than 5% by weight, a effect of added solvent is weak, therefore the transparency electrode having 300 to 900Ω/sq of sheet; resistance and 90% of the transmittance can not be manufactured. A gelatinization is progressed in solution or ununiform film is manufactured, when the amide, sulfoxide or the mixed solvent thereof is more than 25% by weight. Therefore the amide, sulfoxide or the mixed solvent thereof is preferable to use 5% to 25% by weight based; on the total weight of a composition for coating the organic electrode.

Besides, the surfactant and primary alcohol having C1 to C5 are bad for wetting I feature when the transparent polymer substrate such a polyethyleneterepthalate which have a high Patent Application 6 MJM Docket No. 4820-022 Client No. PPW05-076US

surface free energy is coated with the amide, sulfoxide or the mixed solvent thereof. Therefore it is capable of solving the problem to easily form the ununiform layer.

To consider the affinity with the solution, as a primary alcohol, alcohol having C1 to C5 can be used, more preferably isopropanol, ethanol and a methanol is used. When the primary alcohol is less than 5% by weight, it is bad for wetting feature. When the primary alcohol is more than 10% by weight, it is bad for the conductivity. Therefore, it is preferable to use 5% to 10% by weight based on the total weight of the composition for coating the organic electrode.

On the other hand, the surfactant is preferable to be selected at least one of surfactants from the group consisting of a nonjoic surfactant, anionic surfactant, cationic surfactant and a neutral surfactant and B(hydrophilic-lipophilic balance) is within 7 to 20. As a nonionic surfactant, a polyoxyalkylene alkyl ether containing a polyoxyethylene lauryl ether and a polyoxyethylene stearyl ether, a polyoxyalkylene! alkylphenyl ester containing a polyoxyethylene octylphenyl ether and a poly oxyethylene nonylphenyl ether, a sorbitan fatty acid ester containing a sorbitan monolaurate, a sorbitan monostearate and a sorbitan trioleate, a polyoxyalkylene sorbitan fatty acid ester containing a polyoxyethylene sorbitan monolaurate, a poly oxyalkylene fatty acid ester containing a polyethylene monolaurate and a poly oxyethylene monostearate, a glycerine fatty acid ester containing a olenic acid mono glyceride and a stearic acid monoglycerate and a polyoxyethylene-polypropylene block copolymers can be used.

As a anionic surfactant, a fatty acid sodium containing a sodium stearate, sodium oleate and a sodium laurate, a alkylaryl sulfonic acid sodium containing a sodium do decylbezenesulfonate, a alkylsulphuric acid ester sodium containing a sodium lau rylsulfate, a alkylsulfosuccinic acid ester sodium containing a sodium monooctylsul fosuccinic acid, sodium dioctylsulfosuccinate and a sodium polyoxyethylene laurylsul fosuccinate, polyoxyalkylene alkylether sulphuricacidestersodium containing a sodium polyoxyethylenelauryl ethersulfate and a polyoxyalkylene alkylarylether sulphuricacid ester sodium containing a sodium polyoxyethylene nonylphenyl ether sulfate can be used. As a cationic surfactant and neutral surfactant, a alkyl amine sodium containing a; lauryl amine acetate, a 4-level ammonium sodium containing a lauryltrimethylammonium chloride and a alkylbenzyldimethylammonium chloride and a poly oxyethylalkyl-amine can be used. More preferably as a nonione surfactant, a poly oxyethylene surfactant having a excellent wetting feature is used. When the surfactant is less MJM Docket No. 4820-022

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than 0.01% by weight, the form of film is ununiform so that the wetting feature is bad. When the surfactant is more than 0.1% by weight, the surfactant and the nano particle of conductive polymers is phase-separated thereby can be formed nontransparant layer. Therefore the surfactant is preferable to use 0.01% to 0.1% by weight of based on the total weight of the composition for coating the organic electrode.

In case of coating to the composition comprised with the composite, the hardness of the layer of organic conductivity is good, as 3H of pencil hardness tester, but the composition can further comprise the cross-linking agent in order to improve the hardness of layer. As the cross-linking agent which is combining acid group of polystyrenesulfonate(PSS) with hydroxy group of polyhydric alcohol or polyol or is capable of inducing the link with each hydroxy group of the polyhydric alcohol and the polyol, a 4,4-diphenylmethane diisocyanate, toluene diisocyanate, hexamethylene di isocyanate and a organic titanium compound(Vertic IA10, Johnson Mattey Catalysts) can be used. When the cross-linking agent is less than 0.01% by weight, the cross linking is not sufficient and the improvement of the hardness of layer is inadequate. When the cross-linking agent is more than 0.2% by weight, it is difficulty in forming the uniform layer and bad for stability of solution in the long time, because the cross linking agent is tend to gelatinize in mixed solution. Therefore the cross-linking agent is preferably added to use 0.01% to 0.2% by weight of based on the total weight of the composition for coating the organic electrode.

The conductivity of layer feature can be improved by adding a monomer coating a sulfonic acid group(-SO<sub>3</sub>H) as a further dopant to the conductive layer composed the composition. As a dopant, a polystyrene sulfonic acid, p-toluene sulfonic acid, dodecylbenzensulfonic acid, 1,5-anthraquinone disulfonic acid, 2,6-anthraquinone disulfonic acid, anthraquinone disulfonic acid, 4-hydroxybenzensulfonic acid, methylsulfonic acid or a nitrobenzensulfonic acid can be used. When the dopant is less than 0.01% by weight, a dopping effect is decreased. When the dopant is more than 0.5% by weight, the uniformity of the layer is decreased so that added monomer dopant is phase-separated. Therefore, when the dopant is added, the dopant is preferable to use 0.01% to 0.5% by weight to based on the total weight of the composition for coating the organic electrode.

Following, a method of manufacturing organic electrode having excellent transparency using the composition is described in more detail.

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A method of manufacturing organic electrode having excellent transparency according to the present invention comprises steps of stirring the composition for coating the organic electrode, spreading out the stirred composition on the transparent substrate, drying up the substrate and coating to 0.2 to  $20\mu m$  by thickness of coating layer. But according to usage, further comprises steps of repeatedly dispersing the stirred composition 2times to 10 times for 3 to 10 minutes with the ultra sonicator controlled by 20.000 to 40.000Hz of frequency, 50 to 700W of power after the step for stirring.

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First, the composition for coating the organic electrode is manufactured by a order which is slowly stirring the aqueous solution of polyethylenedioxythiophene conductive polymers, at the same time adding orderly the polyhydric alcohol or the polyol, the primary alcohol, amide solvent or the sulfoxide solvent, surfactant, cross linking agent, dopant and then stirring sufficiently at room temperature for 1 to 2time.

In case of manufacturing method using the ultra sonicator, the step of repeatedly I dispersed the composition for 3 to 10 minutes with the ultra sonicator controlled by 20,000 to 40, 000Hz of frequency, 50 to 700W of power is repeated at 2 times to 10 times, so the swelling of conductive nano particle gel is increased by the step. Following, the dispersed solution is spreaded out on the transparent substrate, e.g polyester film, dried up the substrate at heating, so coating layer is formed, the thickness of the coating layer is 0.2 to 20µm, more preferably 0.5 to 10µm. As a transparent substrate, a glass, cellulose ester, polyamide, polycarbonate, polyester, polystyrene, polylolefin, polymetha acrylate, polysulfone, polyethersulfone, polyetherketone, polyetherimide and a polyoxyethylene can be used and more preferably a triacetyl cellulose, polycarbonate or a polyethylene terephthalate is used.

The visible ray transmittance of conductive layer of organic electrode manufactured by the step is more than 90%, the conductivity is generally 300 to  $900\Omega/\text{sq}$ , more preferably less than  $500\Omega/\text{sq}$  and the hardness of layer is the range of 2H to 4H. The transparent organic electrode can be manufactured by means of the method.

To use the method for manufacturing transparency organic electrode, the organic transparency electrode using for various display can be variously manufactured. Also, the organic electrode of the present invention is widely capable of applying to a various field, e.g a electrode or wiring material of organic transistor, smart card, antenna, electrode of battery and Patent Application

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fuel battery, capacitor using for PCB or inductor, closing film of electronic wave, preventing film of static electricity generation and a; sensor as well as the transparent electrode using for display.

### 5 [Mode for the Invention]

The present invention will be described in more detail by way of the following examples, comparative examples, preparative example and example of comparative preparation. However the present invention should not be limited the examples, comparative examples, preparative example and example of comparative preparation.

<Example 1-8>

[Table 1]

		Example (Unit: % by weight)							
		1	2	3	4	5	6	7	8
Conductive	Baytron	remain	remain	remain	remain	remain	remaind	remainder	remainder
polymers	P	der	der	der	der	der	er		
Aqueous	(PEDOT)								
Solution	-								
Polyhydic	Ethylenegl	13	15	13	13	10	13	13	13
alcohol or	ycol(EG)								
polyol									
	Diethylene	-	-	-	-	10	-	-	-
	glycol								
	(DEG)								
Primary	Isopropane	7	9	7	7	7	7	7	7
alchol	ol								!
	(IPA)								
Amide	Methylfor	2	2	2	2	2	15	2	2
Solvent	mamide								

	(MF)								
	Dimethylf ormamide (DMF)	-	-	-	13	-	-		-
	N- Methylpyr rolidone (NMP)	-	-	13	-	-	-	-	-
Sulfoxide Solvent	Dimethyls ulfoxide (DMSO)	13	13	The state of the s	-	-	-	13	13
Surfactant	Triton X- 100	0.06	0.06	0.06	0.06	0.06	0.06	0.06	0.06
Dopant	p- toluenesulf onic acid	-	-	-	-	-	-	0.1	-
	Dodecylbe nznesulfon ic acid		-	-	-	-	-	-	0.1

#### -Preparation-

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As a aqueous solution of polyethylenedioxythiophene(PEDOT) conductive polymers, Baytron P(polyethylene-dioxythiophene;Bayer AG. the location:Leverkusen, Germany) was bottled in a beaker in an amount of the Table 1, and the Baytron P is stirred. As a amide solvent, a methylformamide(Aldrich, the location of Wisconsin, US), dimethylformamide(Aldrich, the location of Wisconsin, US); as a sulfoxide solvent, a dimethylsulfoxide(Aldrich, the location of Milwaukee Wisconsin, US); as a polyhydric alcohol or polyol, a ethyleneglycol(Aldrich, the location of Milwaukee

Wisconsin, US), a diethylene glycol(Aldrich, the location of Milwaukee Wisconsin, US); as a primary alcohol, a isopropaneol(Aldrich, the location of Milwaukee Wisconsin, US); as a Patent Application

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surfactant, a Trion X-100(manufactured by Union carbide); and as a dopant, a p-tolenesulfonic acid(Aldrich, the location of Milwaukee Wisconsin, US) or a dodecylbenzensulfonic acid(Aldrich, the location of Milwaukee Wisconsin, US) in an amount of the table 1 were added into the stirred solution of the Batyron P, therefore the composition for forming film was produced.

<Comparative example 1 ~ 7>

[Table 2]

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		Comparative example (Unit: % by weight)							
		1	2	3	4	5	6	7	
Conductive	Baytron	100	remain	remain	remain	remain	remain	remain	
polymers	P		der	der	der	der	der	der	
Aqueous	(PEDOT)								
Solution									
Polyhydic alcohol	Ethylenegl	-	-	-	-	-	14	16	
or polyol	ycol(EG)								
	Diethylene	-	-	-	10	-	-	-	
	glycol								
	(DEG)								
Primary	Isopropane	-	-	-	-	33	17	4	
alchol	ol								
	(IPA)								
Amide	Methylfor	-	-	3	-	2	-	2	
Solvent	mamide								
	(MF)								
Surfactant	Triton X-	-	0.06	-	-	-	-	-	
	100								

<sup>-</sup> Preparation -

With the exception of the addition in an amount of the table 2, the composition for forming film is produced by the same method with the examples  $1 \sim 8$ .

<Pre><Preparative example 1 ~ 8>

The composition of the example 1 was stirred by 300rpm for 1 time, spreaded out the stirred composition on the polyester film using a Ba coaler, and dried up the film at 100°C dryer for 30minutes, therefore the organic transparent electrode having the form of the transparent substrate and the thickness of coating layer was produced.

<Example 1 of comparative preparation>

As the comparative example 1, the Baytron P was spin-coated on the glass substrate by 300rpm for 30seconds, dried up the substrate at 110°C dryer for 30minutes, therefore the organic transparent electrode having the form of the transparent substrate and 400nm by thickness of coating layer was produced.

<Example 2 of comparative preparation>

With the exception of further addition of a Tritron X-100, the organic transparent electrode was produced by means of the same method with the method of the example 1 of comparative preparation.

<Example 3~7 of comparative preparation>

With the exception of the usage of the composition produced by the comparative example 37 instead of the example 1 of comparative preparation, the organic transparent electrode was produced by means of the same method with method of the example 1 of comparative preparation.

<Experimental example 1>

To use the transparent electrode produced by the preparative example 1~8, the conductivity, transmittance, and the hardness of layer were measured. The conductivity was measured by the sheet resistance with a sheet resistor (Loreasta-GP MCP-T600, Mitsubishi chemical Co.), the transmittance was measured by 550nm transmittance with a UV-vis spectrometer(Helios 0, Spectronic Unicam Co.) and the hardness of layer was measured with a pencil hardness tester, therefore the result by the measurement was shown under the table 3.

30 [Table 3]

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|                             | Preparative example |     |     |     |     |     |     |     |  |
|-----------------------------|---------------------|-----|-----|-----|-----|-----|-----|-----|--|
|                             | 1                   | 2   | 3   | 4   | 5   | 6   | 7   | 8   |  |
| Conductivity( $\Omega$ /sq) | 500                 | 600 | 750 | 700 | 850 | 800 | 400 | 800 |  |
| Transmittance(%)            | 93                  | 90  | 86  | 88  | 86  | 88  | 93  | 92  |  |
| Hardness of layer           | 3H                  | 3H  | 3H  | 3H  | 3H  | 3H  | 3H  | 3H  |  |

As shown in the above table 3, all the transparent electrode produced by the preparation example 1–8 indicated excellence that the conductivity (sheet resistance of layer) is within 300 to  $900\Omega$ /sq, the transmittance of visible ray area is approximate to 90% and the hardness of layer is within 2H to 4H.

## <Experimental example 2>

To use the example 1~7 of comparative preparation instead of the preparative example 1~8, the conductivity, transmittance, and the hardness of layer were measured, therefore the result by the measurement was shown under the table 4.

[Table 4]

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|                             | Example of comparative preparation |                     |     |     |            |     |     |  |
|-----------------------------|------------------------------------|---------------------|-----|-----|------------|-----|-----|--|
|                             | 1                                  | 2                   | 3   | 4   | 5          | 6   | 7   |  |
| Conductivity( $\Omega$ /sq) | 10 <sup>6</sup>                    | 8 x 10 <sup>5</sup> | 600 | 900 | 700        | 700 | 800 |  |
| Transmittance(%)            | 85                                 | 85                  | 70  | 87  | 84         | 83  | 82  |  |
| Hardness of layer           | 2B                                 | 2B                  | 1B  | 1H  | 1 <b>B</b> | 2H  | 2H  |  |

As shown in the above table 4, in the example of comparative preparation 3 and 7, when an amount of aqueous solution of polyethylenedioxythiophene(PEDOT) conductive polymers was more than 90%, the transmittance is not sufficient more than 90%. In the example 1~3 and 5 of comparative preparation, when the polyhydric alcohol was not used, the hardness of layer was decreased and the layer was not uniform.

<Preparative example 9>

The composition of the example 1 was stirred by 300rpm for 1 times, and dispersed the stirred composition with the ultra sonicator controlled by 2,000Hz and 140W, 5times for Patent Application 14 MJM Docket No. 4820-022 Client No. PPW05-076US

3minutes. The dispersed solution was spreaded out on the polyester film with the Barcorder, dried up the film with 110°C dryer for 30minute, therefore the organic transparent electrode having a 2µm by thickness of coating layer was produced.

<Example 8 of comparative preparation>

With the exception of dispersion produced by the preparative example 1, wherein mixture was dispersed by 140W of the ultra sonicator for 10 seconds, therefore the transparent electrode was produced with same method.

<Example 9 of comparative preparation>

With the exception of the produced dispersion by the preparative example 9, wherein mixture was dispersed by 1,000W of the ultra sonicator for 3 minutes, the transparent electrode was produced with same method.

<Experimental example 3>

According to measuring method of the experimental example 1, the conductivity, transmittance and the hardness of layer of the preparative example 9 and the example 8 and 9 of comparative preparation were measured.

[Table 5]

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|                             | Preparative | Example of    | Example of    |
|-----------------------------|-------------|---------------|---------------|
|                             | Example 9   | Comparative   | Comparative   |
|                             |             | Preparation 8 | Preparation 9 |
| Conductivity( $\Omega$ /sq) | 450         | 600           | 104           |
| Transmittance(%)            | 90          | 95            | 88            |
| Hardness of layer           | 3H          | 3H            | 3H            |

As shown in the above table 5, when the preparative example 9 was controlled that the power of the ultra sonicator is the range of 50W to 700W, the dispersion time is 3 to 10 minutes. Therefore the transmittance of visible ray was excellent as 90% and more preferably the conductivity showed  $450\Omega/\text{sq}$ .

#### [Industrial Applicability]

From the above detailed description, a composition for coating an organic electrode and method of manufacturing an organic electrode having an excellent transparency using the composition according to a present invention are capable of producing a flexible transparent organic electrode of large dimension which is excellent to a conductivity and transmittance through steps for coating and printing. Therefore the present invention are capable of increasing the economical efficiency of the process rather than a metal oxide electrode using a existing vacuum process, also is widely capable of applying to a various field of a electrode or a wiring material of organic transistor, smart card, antenna, electrode of battery and fuel battery, a capacitor using for PCB or a inductor, closure of electronic wave and a sensor etc. as well as the transparency electrode using for display.